ZIL'BERMAN. V. M.

USSR/Electricity
Heating, Electric
Electrical Equipment

Apr 1948

THE STATE OF THE S

"Unification of High-Frequency Installations," D. B. Mondrus, S. M. Margolin, V. M. Zil'berman, Engineers, 'ElektroPech'Trust, Ministry of Electrical Industries, USSR, 8 pp.

"Elektrichest" No 4

Soviet industry is sorely in need of invention of a series of apparatus for use in high-frequency heating technology. Mentions work done by SevZapPromElektro-Pech in this field of research and successes achieved. Some machines and equipment have already been placed in various industries and authors ask that workers operating equipment send in testimonials or criticisms.

PA 69T27

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ACCESSION NR: AP4035099

s/0191/64/000/005/0013/0015

AUTHOR: Smirnova, O. V.; Losev, I. P. (Deceased); Yerofeyeva, S. B.; Zil'berman, Ye. G.

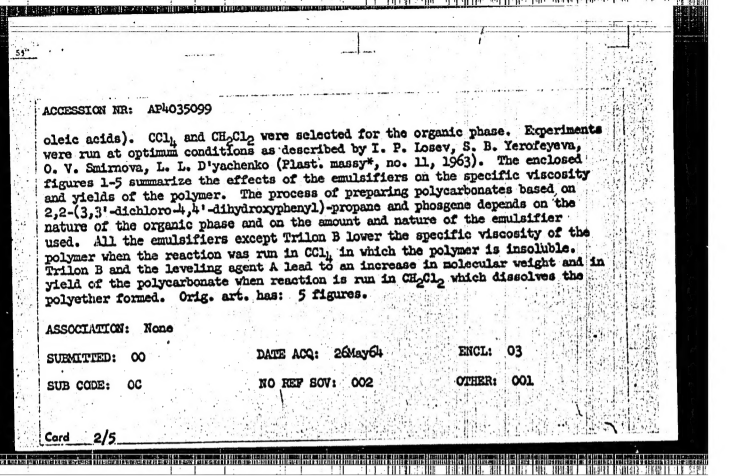
TITIE: Effect of emulsifiers on the course of the interphase polycondensation reaction in preparing polycarbonates based on dichlorodiphenylolpropane.

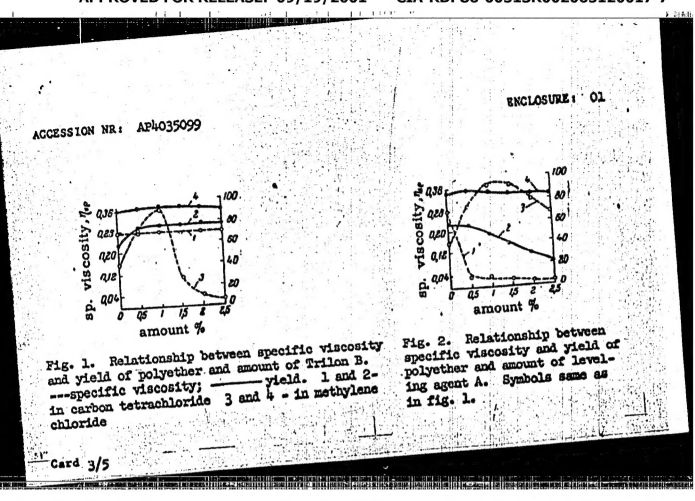
SOURCE: Plasticheskiye massy*, no. 5, 1964, 13-15

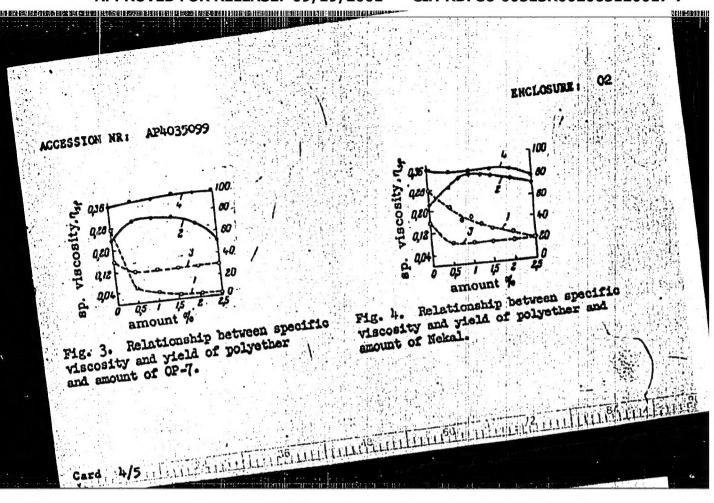
TOPIC TAGS: emulsifier, interphase polycondensation, polycarbonate, dichlorodiphenylolpropane, dichlorodiphenylolpropane polycarbonate, Trilon B, leveling agent A, OP 7, Nekal, Avirol, molecular weight, yield, specific viscosity, polyether, solvent effect

ABSTRACT: The effect of certain emulsifiers on the molecular weight and the yield of polycarbonates based on a chlorinated dihydroxydiphenylalkane were investigated. The following emulsifiers were studied: Trilon B, leveling agent A (quaternary ammonium salt of diethylaminomethyl derivatives of polyethylane glycol esters of isooctylphenols), OP-7 (polyethylane glycol ester of isooctylphenol), Nekal, and Avirol (mixture of neutral esters of butanol, sulfuric and

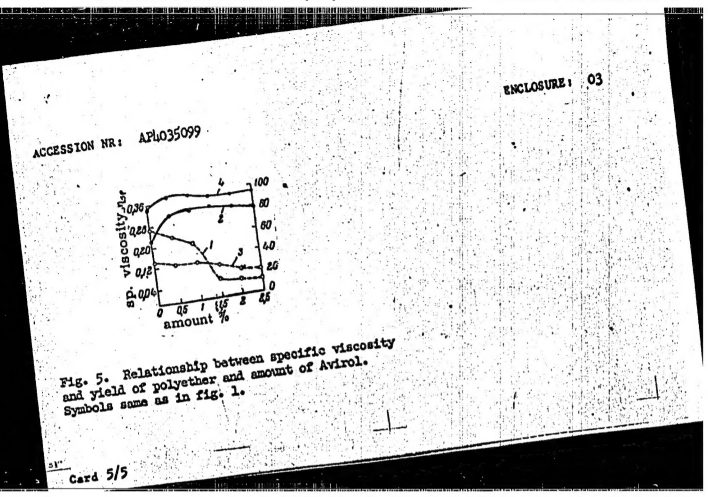
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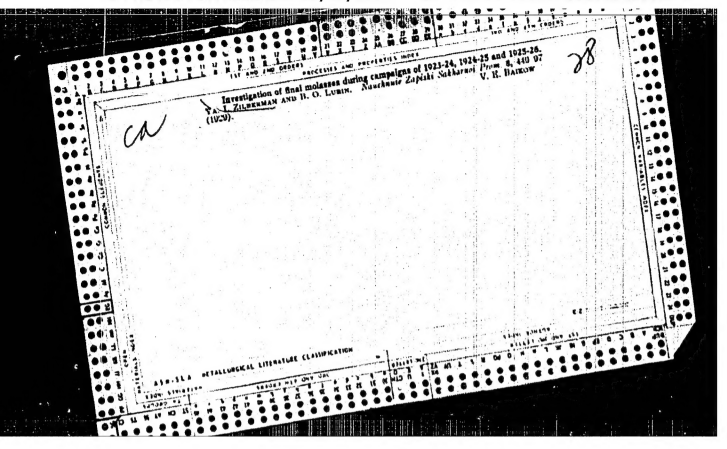


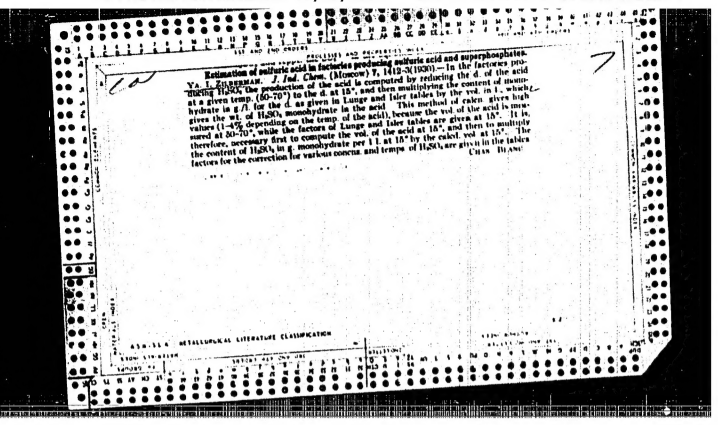


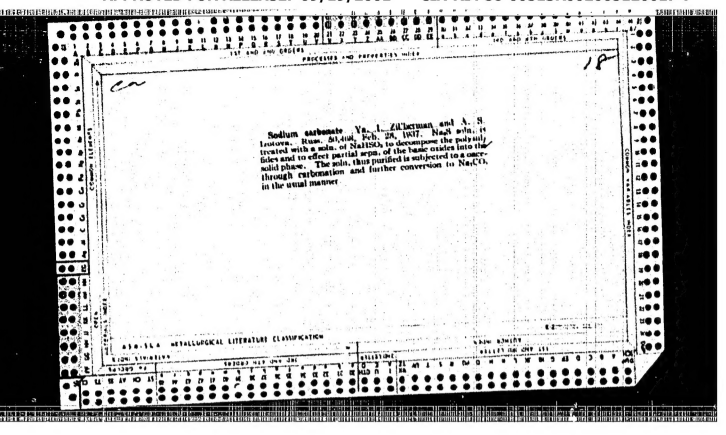
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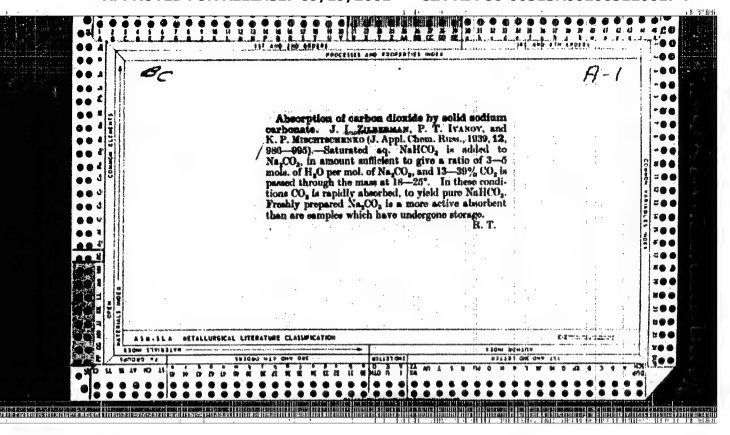


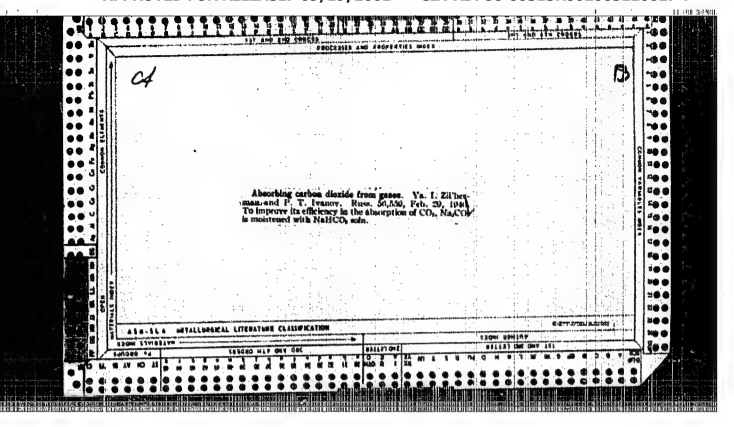
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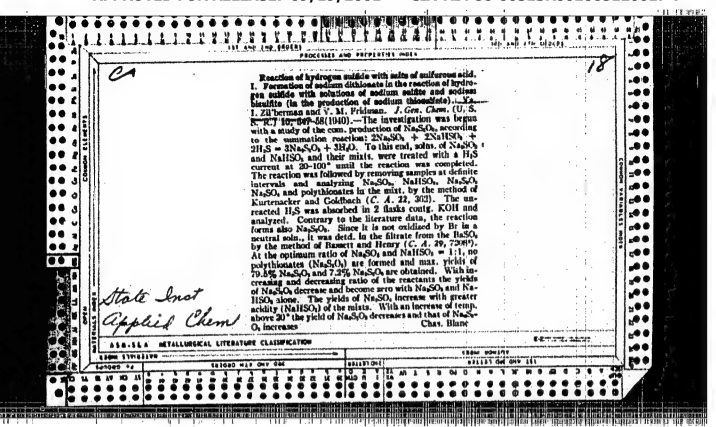


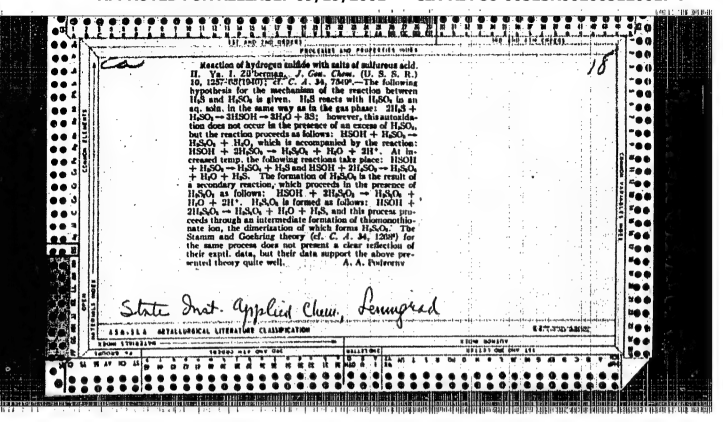


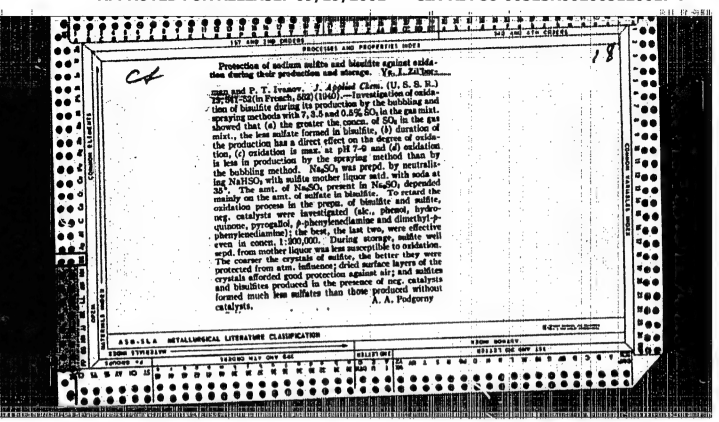


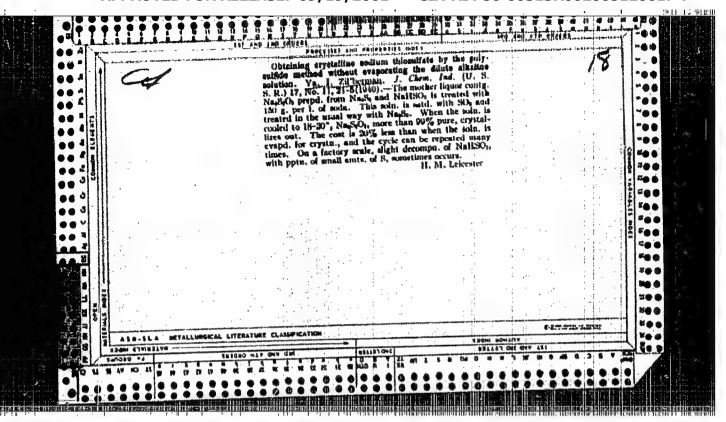


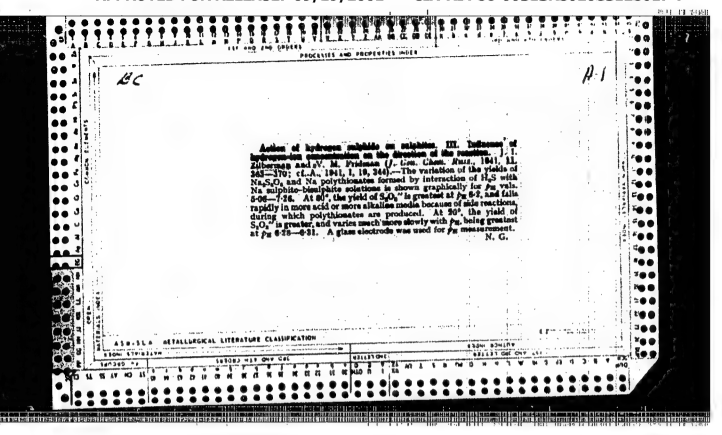


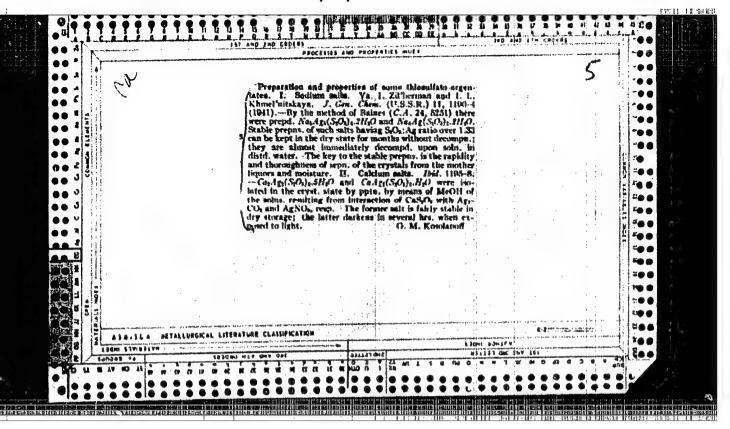


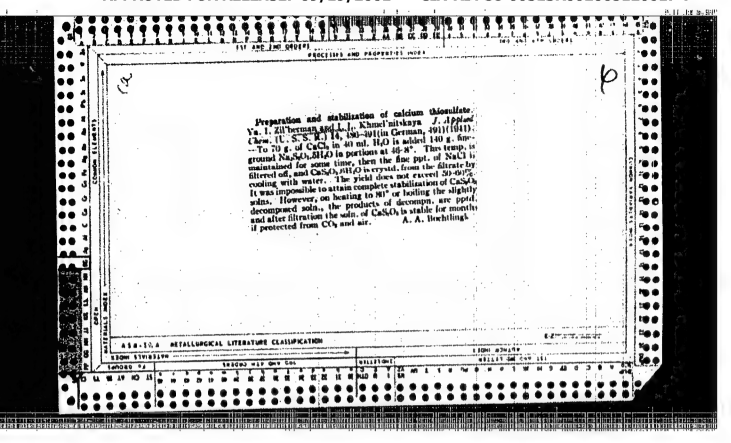


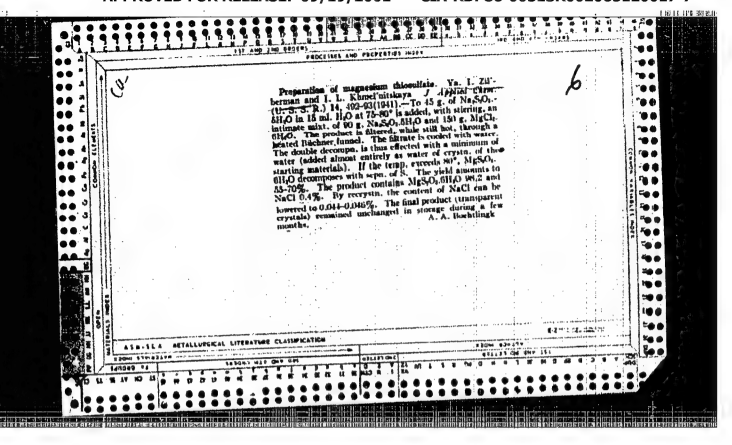


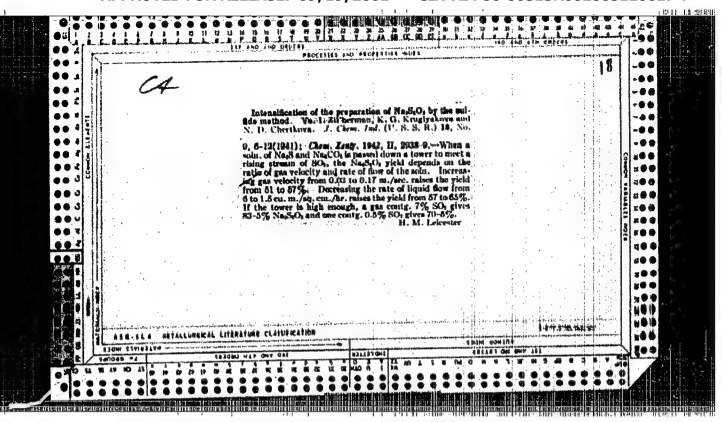


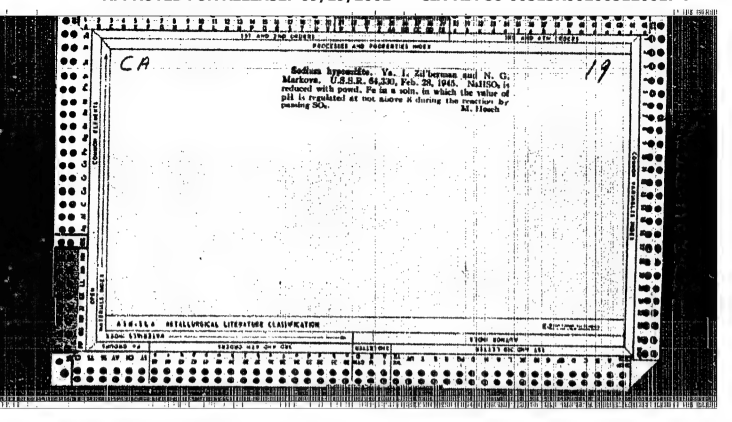


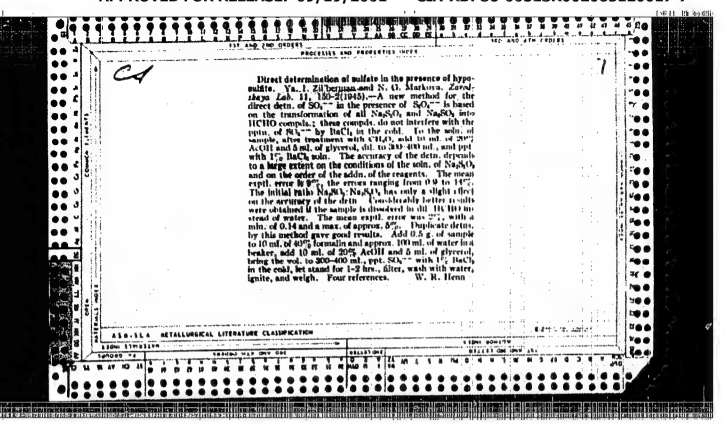


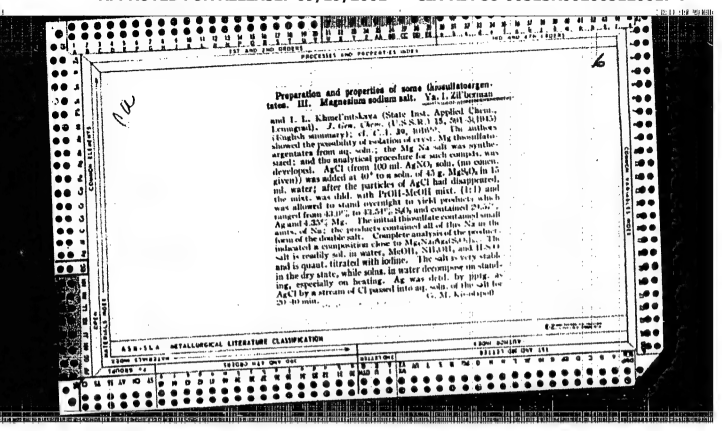


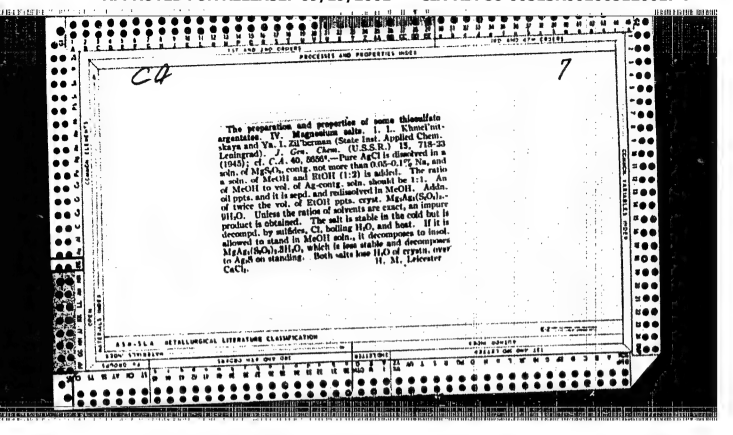


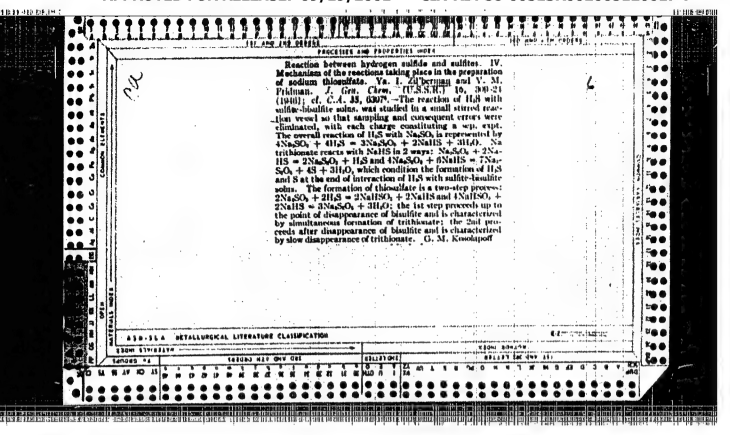


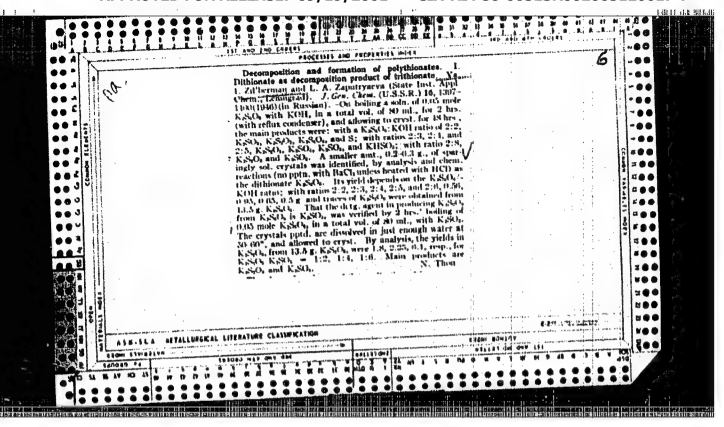


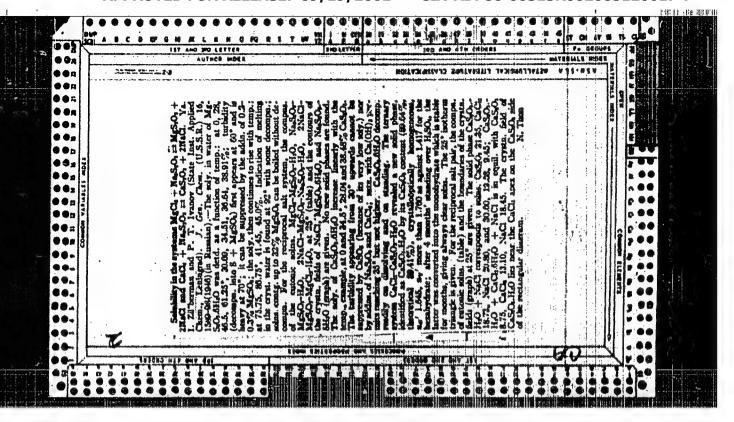


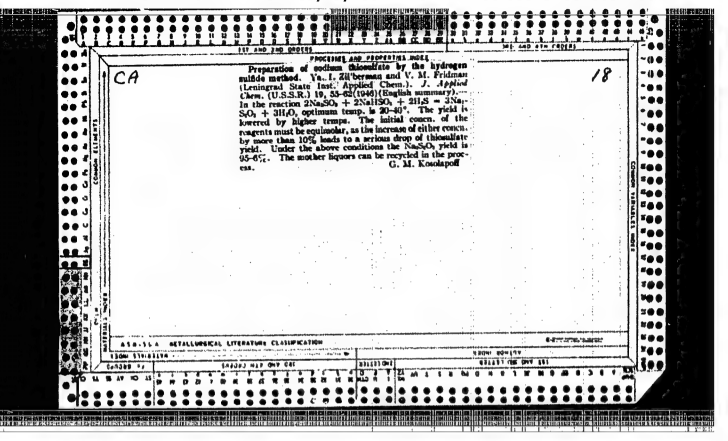


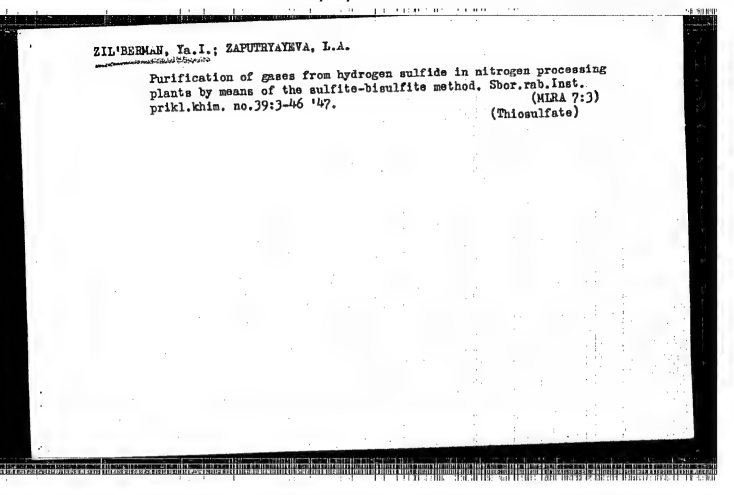


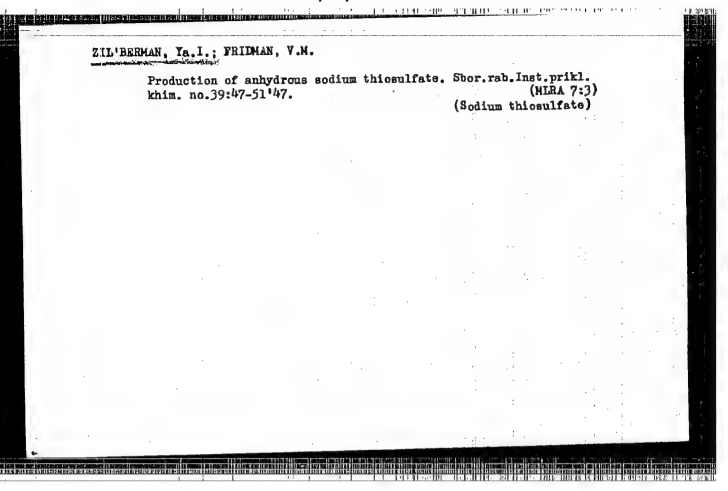


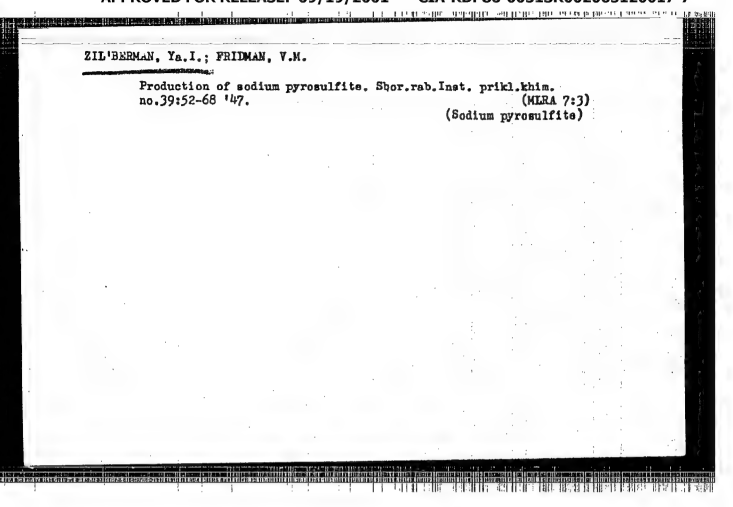












8/186/60/002/006/005/026 A051/A129

AUTHORS:

Zil'berman, Ya. I.; Peshchevitskiy, B. I.

TITLE:

The production, composition and certain properties of uranyl

disthyldithiocarbamate complexes

PERIODICAL:

Radiokhimiya, v. 2, no. 6, 1960, 663 - 667

TEXT: The article deals with a description of a method for producing salts of the general formula Mⁿ⁺ (UO₂Car₃)_n, where Mⁿ⁺ is the cation of alkaline or alkaline-earth metals, ammonium or distinglarmonium and Car is the diethyldithicoarbamate ion. The properties of these salts are discussed. In addition, the uranyl tri-carbamates with rubidium, cashum calcium, strontium, barium and ammonium were produced for the first time in the external sphere of the complex. The properties of ten different diethyldithicoarbamate derivatives of uranyl have been characterized. During the experiments the electroconductivity was measured on an apparatus consisting of a sound generator, bridge, and -lectron-heam indicator of equilibrium. The optical density of the solutions was measured on a two-arm \$\dagger{q}\$EM (FEX-M) photocolorimeter. The method of analysis of the metal con-

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The production, composition and

tent enabled the authors to show that the residue obtained after decomposition and calcination of the complexes is a diuranate of the corresponding metal. In the case of the dicarbanate complex and the ammonium derivatives the expected uranous-uranic oxide is obtained. When heating to 11000 a darkening of color and loss of weight was noted in some compounds. In moist atmosphere the salts regained their initial properties which points to the presence of crystallization water in the complexes. A good coincidence of experimental and theoretical values was observed. The dry, pure complexes are found to have a high thermal stability. All the frashly prepared compounds are soluble in water, the "aging" affect is observed, however, when these are stored. A drop in the solubility is noted at an increase in the ion radius of the one-charge cation of the external sphere (from sodium to essium) The complexes obtained are soluble in many organic solvents and can be recrystallized from the latter. The organic solvents follow the sequence of solubility: ketones = alcohols > esters = chloroderivatives > ethers ~ aromatic hydrocarbons, whereby the latter two classes can only dissolve uranyl dicarbamate. Aqueous solutions of uranyl carbamates are decomposed by acids, alkalies and soda solutions. In an acid medium the stability depends to a great extent on the pH of the solution and a noticeable change in

Card 2/3

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The production, contestition and

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the stability begins at pH > 4.5. Senerally speaking, the stability of the complexes surpasses that of the non-bound dithiocarbanic acid. (Ref. 12: H. Bode Z. Anal. Chem., 142, 6, 4:4, 1954). The molecular slectroconductivity of the sodium uranyleschamate solutions was found to be equal to about 95 chm⁻¹. cm⁻¹, 1.e., corresponding to 1: 1 of the electrolyte. In conclusion the authors point out that the quantitative results of the analyses on solubility should be regarded with respect to the aging phenomenon, even though these results give a general picture of the solubility properties. There are 3 tables, it figure and 12 references: 5 Soviet-bloc and 7 non-Soviet-bloc. The references to the English language publications rate as follows: P. I. Lancete, M.S. Earing, Anal. Chem., 23, 6, 871., 1951. R. A. Zingaro, S. Am. Chem. Soc., 78, 15, 3518, 1956.

SUBMITTED:

Department 10, 1989

Card 3/3

PHASE I BOOK EXPLOITATION SOV/5448

Zil'berman, Ya.I.

Osnovy khimicheskoy tekhnologii iskusstvennykh radioaktivnykh elementov (Chemical Technology Principles of Artificial Radioactive Elements) Moscow, Gosatomizdat, 1961. 331 p. Errata slip inserted. 6,000 copies printed.

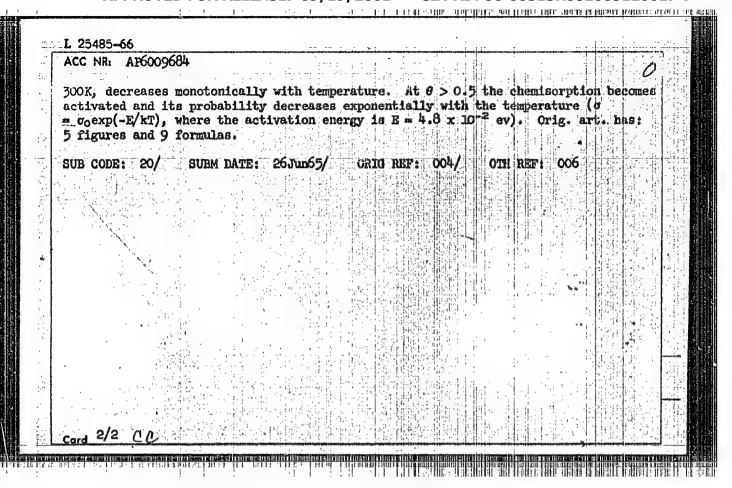
Ed.: Ye.I. Panasenkova; Tech. Ed.: N.A. Vlasova.

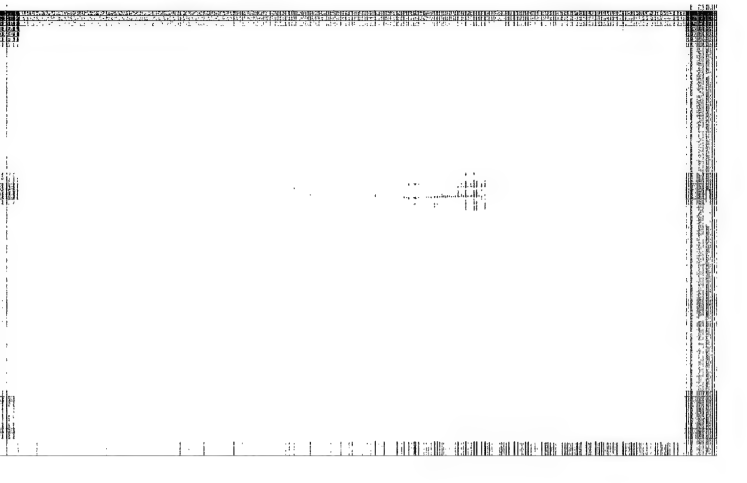
PURPOSE: This book is intended for radiation chemists and other specialists and scientific workers interested in the production of artificial radioactive elements.

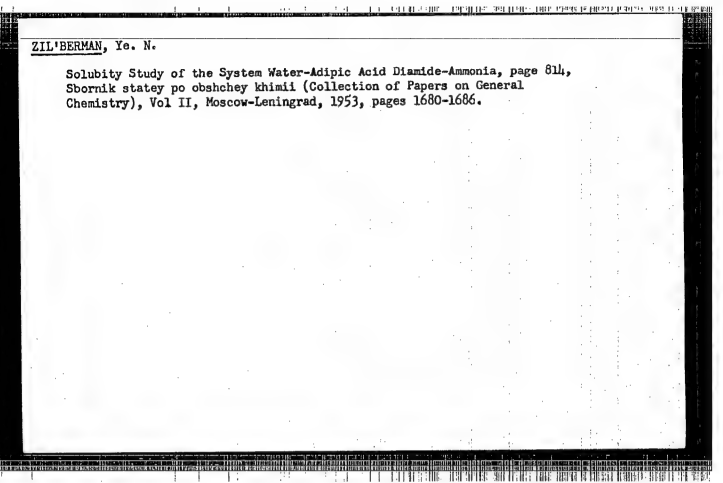
COVERAGE: The book gives a systematic presentation of data on the chemical technology of artificial radioactive elements, especially nuclear fuel processing. Methods for processing irradiated material are discussed, as are the biological and chemical effects of ionizing radiation. Methods for nuclear fuel processing which are still in a stage of development at laboratories and experimental plants are not dealt with. Theoretical principles are treated only to the extent

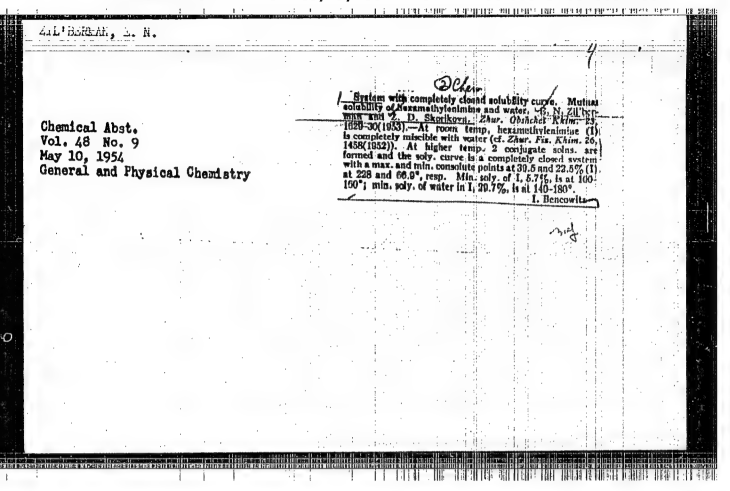
Card 1/7

ACC NRi AP6009684	80URCE CODE: UR/0181/66/008/003/0912/0919
AUTHOR: Zingerman, Ya. P.; Ishchuk,	
ORG: Institute of Physics; AN Ukrss	R, Kiev (Institut fizili AN UkrSSR)
TITLE: Adsorption of oxygen on the	(110) face of single-crystal tungsten
SOURCE: Fizika tverdogo tela, v. 8,	no. 3, 1966, 912-919 (8 2
TOPIC TAGS: tungsten, single crysts sorption	l, gas adsorption, oxygen, crystal surface, chemi-
ABSTRACT: This is a continuation of	earlier work by the authors (FTT v. 7, 2569, between oxygen and surfaces of different faces of
tungsten single crystals. The prese	ent investigation was devoted to the (110) face,
and the experimental equipment used	was the same as in earlier studies. The method blecular oxygen from a beam under conditions of
very high vacuum (~10-10 mm Hg). The	ne single crystal was grown by zone melting. The
results show that the interaction wi	th the (110) face consists only of adsorption of sociation and formation of a final structure of
the WO type. It was observed that t	the (110) face is characterized by the presence of
only one type of oxygen chemisorptic	on state with the binding energy between the ad- ev, regardless of the degree of coating of the
surface 6, with the latter ranging i	from $\theta \le 0.1$ to $\theta \simeq 0.9$. In the region $\theta \le 0.5$,
the chemisorption is not an activate	ed process, and its probability, equal to 0.1 at
Card 1/2	









ZIL'HERMAN, Ye.N.				. • []		
Ac 153.		and water.	Zhur.prikl.khim. 26 no.8:886-88 (MLRa 6:8) (Azeotropy) (Cyclohexanol)			
(CA 47 no.22:11)	100, 600		(22000100) (0302	Green Town		
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	System: water	r - adiponitrile - ammonia. Zhur.prikl.khim. 26 no.9:941-940 (MIRA 6:10						948 6=10)))	
		(Systems	(Chemistry))	(Adiponitr	lle)	(Ammon	ia)			
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and a second of the light part and the first of the control of the second secon ZIL'BERMAN, 10. A. USSR/Chemistry - Production of Cyclohexanone FD 169 Card 1/1 Author : Zil'berman, Ye. N. : Thermodynamic investigation of the processes for the production of cyclo-Title hexanol and cyclohexanone from phenol. Periodical: Khim. prom. 3, 24-26 (152-154), April-May 1954 On the basis of thermodynamic consideration, arrives at the conclusion Abstract that the almost quantitative yield of cyclohexanone which is supposed to result from the interaction of phenol with cyclohexanol according to British Patent 310055 is improbable. States that in the presence of suitable catalysts this dismutation may proceed to the extent of 50% per run, however, so that the process is industrially feasible. 8 USSR references and 11 foreign references are appended.

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AID P - 3755

Subject

: USSR/Chemistry

Card 1/1

Pub. 152 - 19/22

Authors

Title

Zil'berman, Ye. N. and G. N. Matveyeva

The control of the co Products resulting from the interaction of hexamethylene diamine with some inorganic acids

Periodical

Zhur. prikl. khim. 28, 9, 1013-1016, 1955

Abstract

Mixtures of basic salts with neutral salts and free hexamethylene diamine were obtained in the reactions of hexamethylene diamine with sulfuric and with nitric

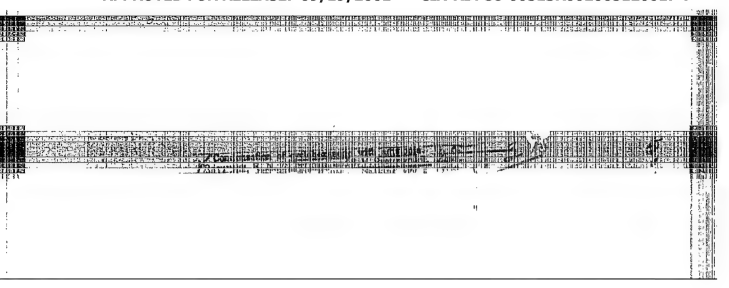
acids. Eleven references, 2 Russian (1947-1953).

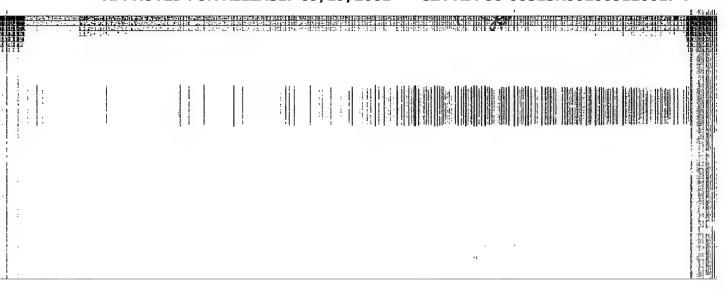
Institution :

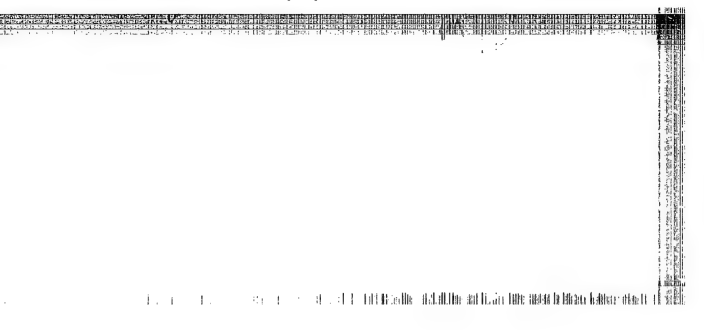
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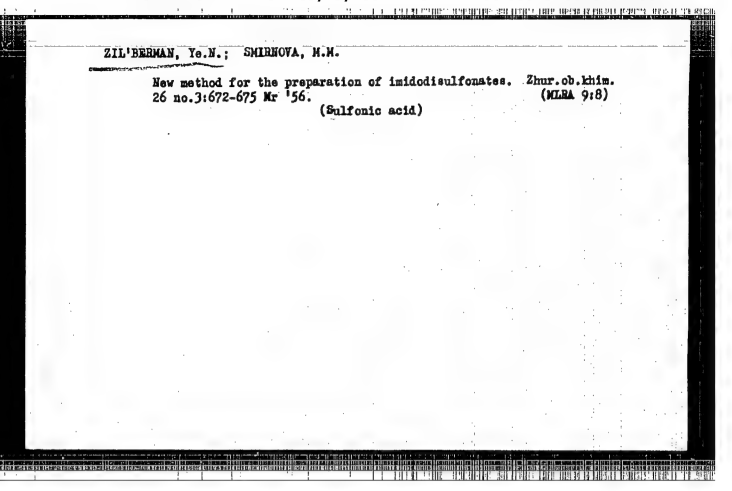
Submitted

: Mr 19, 1954









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USSR/ Organic Chemistry - Synthetic organic chemistry

Abs Jour : Referat Zhur - Khimiya, No 4, 1957, 11632

Author

Zil'berman Ye. N., Suvorova S.N., Smolyan Z.S.

Title

: On Preparation of Adipic Acid by Oxidation of Cyclochexanol with

Nitric Acid

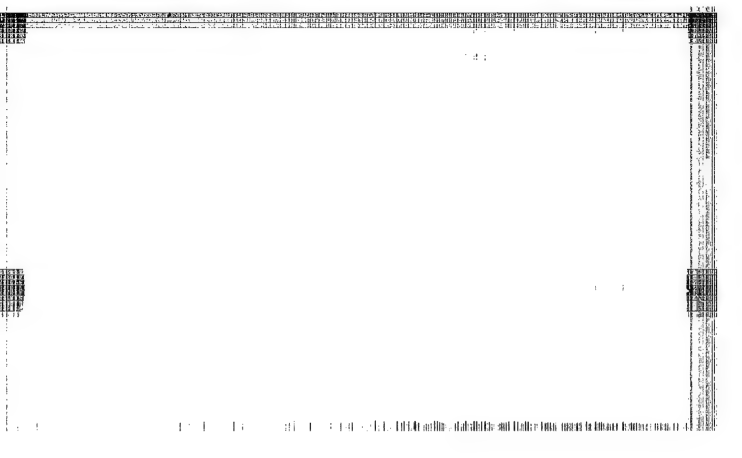
Orig Pub : Zh. prikl. khimii, 1956, 29, No 4, 621-627

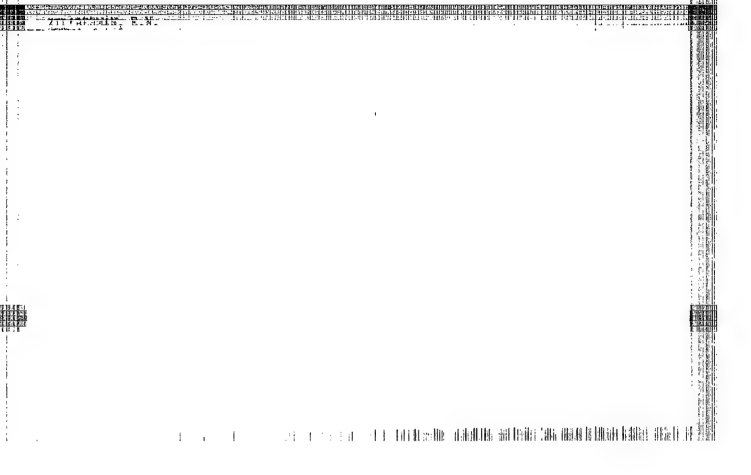
Abstract : Studied was the effect upon the reaction of oxidation of cyclohexanol (I), by the action of nitric acid, to adipic acid (II), of catalysts (CT), duration of reaction, concentration of HNO₂ and the presence therein of organic acids. Maximum yield of II 81-83%; minimum yield of by-products: glutaric (III) (5.6%), succinic (IV) (3.9%), oxalic (V) (4.2%) acids, is obtained with HNO₂ concentration of 40-50%. With decreasing concentration of HNO₃ yield of II decreases and that of III and IV increases; with 5-10% HNO3 the main reaction product is III. In presence of CT (NH4VO3, CuCO3, Bi(NO3)3.6H2O and NH4VO3 +

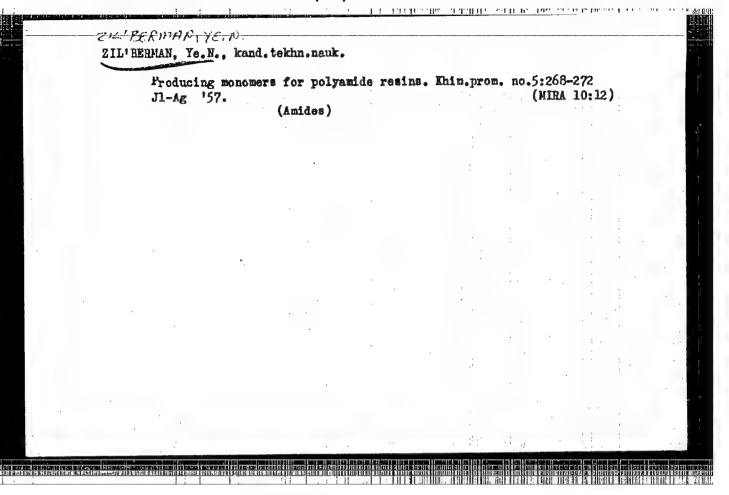
CuCO₂ (1:3) yield of II increases and that of IV decreases. NH_LVO₂

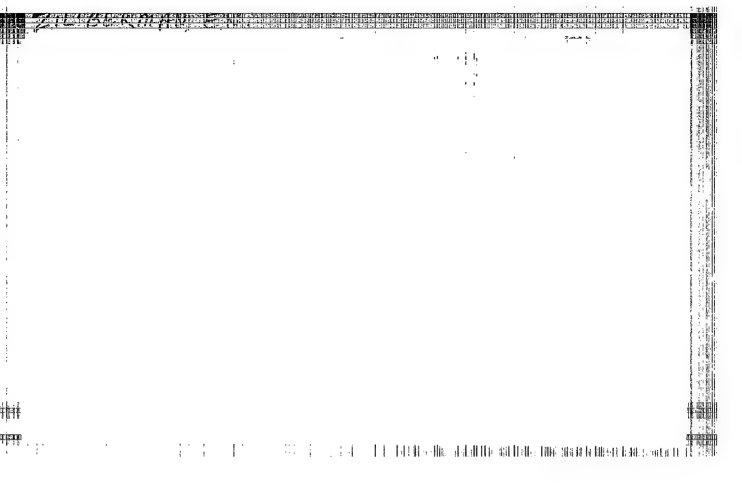
inhibites formation of V. In the presence of CuCO3 formation of III

Card 1/2









Zil' berMAN, ye.N.

AUTHORS:

Zil'berman, Ye.N., Smirnova M.M.,

20-5-21/54

atem the contribute of the technic contribution of the

TITLE:

The Mechanism Underlying the Oximation of Cyclohexanone by Sodium Hydroxylaminmonosulphonates (Abbrev. HAS) (O mekhanizme oksimirovaniya tsiklogeksanona gidroksilaminmonosul'fonatom

natriya)

PERIODICAL:

Doklady Akad. Nauk SSSR, 1957, Vol. 115, Nr 5, pp. 927-930, (USSR)

ABSTRACT:

For the eximation of aldehydes and ketons usually salts of hydrosylamin are used. The industry, however, uses sometimes for the purpose described in the title the salt named last in the title. It is and intermediate product of the synthesis of hydroxylaminsulphate(according to Raschig). The mechanism dealt with here has not been described inscientific periodicals. The following characteristics were found in the reciprocal action of the 2 substances, named in the title: 1) Unlike the hydroxylamin the substance HAS, named last, does not react to the alcaline milieu. 2) This substance HAS does not hydrolize at room temperature, which is at the same time the temperature prevailing at the experiment. 3) With the presence of cyclohexanon and an acid HAS reacts easily, forming cyclohesanonoxim and hydroxylamin. For better understanding the reaction should be written down. Here carboniumion is the primary reaction product. In the further course it establishes a binding of coordinations on account of the unseperated electron pair of the nitrogen atom within the nucleophil

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20-5-21/54

The Mechanism Underlying the Oximation of Cyclohexanone by Sodium Hydroxylaminmonosulphonates (Abbrev. HAS)

stant. Description of experiments with the usual information follow. There are 2 figures, † table and † Slavic reference.

PRESENTED

By Kazanskiy, B.A., Academician, May 15, 1957

SUBMITTED

July 19, 1956

AVAILABLE

Library of Congress.

CARD 3/3

THE PROPERTY OF THE PROPERTY O

SMIRNOVA, O. V.; LOSEV, I. P.[doceased]; YEROFEYEVA, S. B.; ZIL BERMAN, Ye. G.

Effect of emulsifiers on the course of the interphase polycondensation in the production of polycarbonates based on dichlorodiphenylol propane. Plast. massy no. 5:13-15 164.

(MIRA 17:5)

Interaction of acid sulfate ester of A-hydroxypropionamide with alcohols. Zhur. ob. khim. 33 no.8:2766-2771 Ag '63.

(MIRA 16:11)

1. Gor'kovakin politekhnicheskiy institut.

MICHURIN, A.A.; ZIL'HERMAN, Ye.N.

Synthesis of N,N'-diaryl-P-aminopropionamides. Zhur.ob.khim. 34 no.2:575-579 F '64. (MIRA 17:3)

1. Gor'kovskiy politekhnicheskiy institut.

807/64-58-6-13/15 Zil'berman, Ye. N., Fedoseyeva, G. T.

On the Separation of Adiponitrile Obtained From Products of AUTHORS:

the Reaction Between Adipic Acid and Ammonia (O vydelenii adiponitrila iz produktov vzaimodeystviya adipinovoy kisloty

Khimicheskaya promyshlennost', 1958, Nr 6, pp 377-379 (USSR)

One of the methods used for the industrial production of adipo-PERIODICAL: ABSTRACT:

nitrile is the synthesis obtained from adipic acid and ammonia producing a heterogeneous mixture (Ref 1). Adiponitrile obtained in such a way is mostly dissolved in water. It is stated that the aromatic carbohydrates are a good extracting agent for adiponitrile since they do not react with it and have a relatively high steam pressure and specific weight. In order to study these possibilities of extraction analyses were, in the present case, carried out with the systems wateradiponitrile-benzene, water-adiponitrile-toluene, and wateradiponitrile-(NH₄)₂SO₄. Since good results can also be obtained by salting out the aqueous adiponitrile solutions, the

analyses were carried out in the latter system. In a few cases the water content of the oil layer was determined by means of

Card 1/2

TITLE:

On the Separation of Adiponitrile Obtained From Products of the Reaction Between Adipic Acid and Ammonia

the Fischer reagent (Ref 10). The comparison of the extractability of some carbohydrates is illustrated graphically on the basis of the data of the distribution of adiponitrile between water and the solvent. It appears from this that the introduction of a methyl-substitute into the benzene-ring reduces the concentration ratio of adiponitrile between the oil and the aqueous layer. The extractability decreases in the following order: benzene, toluene, ortho-xylene. On the basis of the observations made it is assumed that an extraction in the presence of inorganic salts would be especially effective. There are 5 figures, 1 table, and 10 references, 4 of which

Card 2/2

AUTHORS:

Roginskaya, Ts. N., Svetozarskiy, S. V., 507/79-28-3-47/66

Finkel'shteyn, A. I., Zil'berman, Yc. N.

TITLE:

Concerning the Question of the Molecular Structure of the Unsaturated Ketones Which Are the Bimolecular Condensation Product of Cyclohexanone (K voprosu o molekulyarnom stroyenii nenasyshchennykh ketonov-bimolekulyarnykh produktov konden-

satsii tsiklogeksanona)

PERIODICAL:

Zhurnal obshchey khimii, 1958, Vol. 28, Nr 8,

pp. 2229 - 2233 (USSR)

ABSTRACT:

On the basis of investigations on the chemical properties of the ketone C₁₂H₁₈O(I) the structure (A) (Refs 6,9-11) or structure (B)(Refs 9,10,12-12) may be assigned to it, or it may be considered as a mixture of the two isomers (Reis 15,16). Those supporting structure (B) (Refs 10,14) base their arguments on the chemical reactions of the ketone and cite the rule of Dikman-Kon(Dikman, Kon), according to which the semicyclic double bond in the

Card 1/3

cyclohexane ring is less stable than the endocyclic double bond. The question of the structure of this ketone was more

Concerning the Question of the Molecular Structure of SOV/79-28-8-47/66 the Unsaturated Ketones Which Are the Bimolecular Condensation Product of Cyclohexanone

disputed by the discovery of a new unsaturated ketone C₁₂H₁₈O(II), by Reese (Rize)(Ref 12) in 1942. According to his reactions there could be not doubt that α, β double bonds were present. In contrast to (I) this compound is a solid, relatively less stable, and on warming changes to the liquid ketone (I). The spectra of these two ketones, (I) and (II), had not previously been investigated. The authors investigated the optical properties of (I) and (II) in order to establish their molecular structures. The combined spectra obtained are given in the table, while the infra-red absorption spectra appear in figures 1 and 2: and the ultra-violet spectra appear in figures 3 and 4. It was shown that both compounds are different forms of the α, β -unsaturated ketone 2-cyclohexylide cyclohexane. Figures 1-4 illustrate the spectral analytical results; figure 5 shows the structure of the two stereoisomers of 2-cyclohexylide cyclohexane. There are 5 figures, 1 table, and 20 references, 2 of which are Soviet.

Card 2/3

Concerning the Question of the Molecular Structure of SOV/79-28-8-47/66 the Unsaturated Ketones Which Are the Bimolecular Condensation Product

SUBMITTED: May 22, 1957

Card 3/3

SOY/80-59-1-40/44

AUTHORS:

Kalikova, A.Ye., Zil'berman, Ye.K., Roginskaya, Ts.K. and

Smirnova, M.M.

TITLE

On Purifying Adipo-Nitryl (Ob ochistka adiponitrila)

PERIODICAL:

Zhurnal prikladnoy khimii, 1959, Kr 1, pp 227-230 (USSR)

ARDURACT:

Adiponitryl is an intermediate product in the industrial synthesis of the adipic hexamethylenediamine which is used in the production of polyamide resins. The usual methods employed in this synthesis do not ensure the wanted purity of the adiponitryl. In the present notice the authors suggest, on the basis of chemical analyses and studying the ultraviolet spectrum of the adiponitryl, a new method of its purifying. They show that by treating adiponitryl with sulfuric acid and its subsequent flushing with ammonium bisulfite solution it is possible to obtain the pure and stable (in storing) product which practically does not absorb ultraviolet rays in the range from 220 to 400 mm

Card 1/2

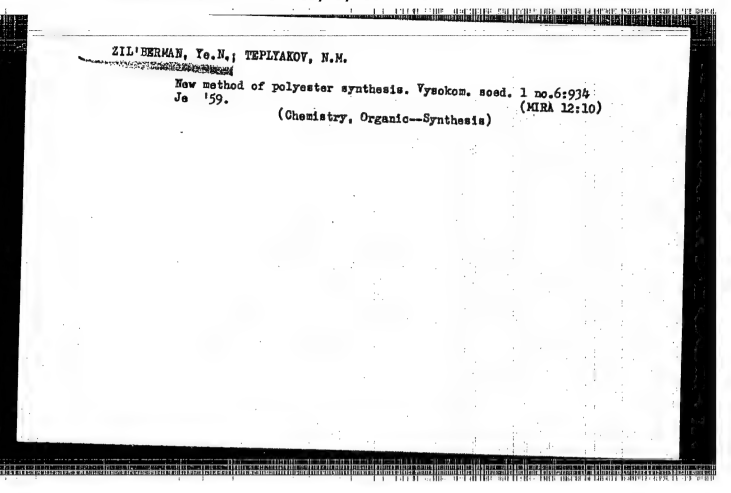
wavelength.

On Purifying Adipo-Nitryl:

There are 4 graphs and 15 references, 4 of which are Soviet, 5 American, 3 German, 2 French and 1 English.

SUBMITTED: May 16, 1957.

Cara 2/2



5(3)

SOV/63-4-1-26/31

AUTHORS:

Zil'berman, Ye.N., Kulikova, A.Ye., Sazanova, N.A.

TITLE:

Method for Preparing Amides From Nitriles (Sposob polucheniya

amidov iz nitrilov)

PERIODICAL:

Khimicheskaya nauka i promyshlennost', 1959, Vol 4, Nr 1,

pp 135-136 (USSR)

ABSTRACT:

A convenient method for the transformation of nitriles to amides in the cold and in the presence of hydrogen chloride is shown here. The interaction product of 1 mole of benzonitrile and two moles of hydrogen chloride is treated with 1 mole of water which produces the compound C6H5CONH2 · HC1. On dissolving it in water it is transformed into benzamide. The hydration reactions of nitriles proceed not only in ether, but also in other organic solvents, like dioxane, benzene, CCl4, etc.

SUBMITTED:

There are 2 references, 1 of which is Soviet and 1 German. July 21, 1958

Card 1/1

5(4) AUTHORS:

Svetozarskiy, S. V., Zillberman, Ye. N., Razuvayev, G. A.

TITLE:

Low-Temperature Autocondensation of Cyclohexanone (Nizkotempera-

turnaya avtokondensatsiya tsiklogeksanona)

PERIODICAL:

Zhurnal obshchev khimii, 1959, Vol 29, Nr 5,

pp 1454 - 1457 ω(USSR)

ABSTRACT:

In the present paper a C₁₈H₃₀O₃ compound (I) was obtained by means of autocondensation of cyclohexanone at room temperature in the presence of solid sodium hydroxide as catalyst. It represents a tricyclic product with a carbonyl- and two ternary hydroxyl-groups. On heating with solid sodium hydroxide (I) decomposes to give cyclohexanone and 3-cyclohexylidene—cyclohexanone. Owing to the reaction of (I) with concentrated sulfuric acid as well as on short heating 2 water molecules are splitted off and an unsaturated C₁₈H₂₆O-ketone (II) is formed.

On protracted heating of (I) and (II) the dodecahydro-

Card 1/2

1,2,3,4,5,6,7,8,9,10,11,12-triphenylene (III) is formed in good yield. Owing to the transformation of (I) into (III) the com-

CIA-RDP86-00513R002065120017-7

Low-Temperature Autocondensation of Cyclohexanone

SOV/79-29-5-10/75

pound C₁₈H₃₀O₃ must be a dioxy-ketone with the structure of the 2-[2-(1-oxy-cyclohexyl)-1-oxy-cyclohexyl]-cyclohexanone (Fig 1). The easy transformation of (II) into (III) may be due to steric factors. In order to define the structure of the ketone (II) its ultraviolet spectrum in diethyl ester (Fig 2) was taken. It was found that (II) can only have the structure of 2-[2-(\alpha'-cyclohexenyl)-cyclohexylidene]- cyclohexanone. No usual derivatives of carbonyl compounds could be obtained from the ketone (II) and the dioxy-ketone (I) which is due to steric hindrances. It has to be mentioned that (III) is usually obtained in a yield of 6% at the most (Ref 3). The transformations of (I) and (II) into (III) here described are a new and convenient method for the preparation of dodecahydrotriphenylene. The authors express their gratitude to Ts. N. Roginskaya for taking the ultraviolet spectra. There are 3 figures and 9 references, 1 of which is Soviet.

SUBMITTED:

April 11, 1958

Card 2/2

5(3) AUTHORS:

SOV/79-29-5-61/75 Zil'berman, Ye. N., Kulikova. A. Ye.

TITLE:

Products of the Reaction of Adiponitrile With Hydrogen Chloride and Their Hydrolysis (Produkty vzaimodeystviya adiponitrila

s khloristym vodorodom i ikh gidroliz)

PERIODICAL:

Zhurnal obshchey khimii, 1959, Vol 29, Nr 5, pp 1694-1699

(USSR)

ABSTRACT:

In the hydrochlorination of adiponitrile two molecules of hydrogen chloride first combine selectively, with a nitrile group and the hydrochloride of imonium chloride of the 6-cyanovalerianic acid is formed. A further hydrochlorination yields dihydrochloride of diimonium chloride of adipinic acid. Chlorides of imonium hydrin passing over to the amines of the corresponding acids by hydrolysis and neutralization form quantitatively by the reaction of equivalent quantities of hydrochlorides of imonium chlorides and water. A new method is suggested for the hydrolysis of nitriles in the presence of concentrated halogen hydracids. New methods of producing δ-cyanovaleramide, δ-cyanovalerianic acid and adipamide in

Card 1/2

high yields were introduced. K. K. Ish cooperated in the experimental work. There are 2 tables and 12 references,

Products of the Reaction of Adiponitrile With Hydrogen Chloride and Their Hydrolysis

4 of which are Soviet.

SUBMITTED: March 17, 1958

Card 2/2

5(3) AUTHORS:

Zil'berman, Ye. N., Kulikeva, A. Ye.

507/79-29-9-50/76

TITLE:

Synthesis of the Imino Esters and the Esters of &-Cyanovaleric

Acid

PERIODICAL:

Zhurnal obshchey khimii, 1959, 7cl 29, Nr 9, pp 3039-3041

(USSR)

ABSTRACT:

At present it is generally assumed (Refs 1-5) that an equivalent amount of nitrile, alsohol, and hydrogen halide is necessary for the formation of imino esters and that the reaction takes place in a stage according to the pattern

RCN + ROH + HCl ->RC NH. HCl. In this case, however, the

yields are low and the final products are rarely obtained in pure state (Refs 4.5). In continuation of a preceding paper of the authors (Ref 6) and on the basis of the publication by I. Guben (Ref 4) on the participation of two molecules of hydrogen chloride in the formation of an imino ester group which is not quite clear it was assumed to be more expedient to carry out the synthesis of the imino esters in two stages, i.e. to add first two molecules RC1 to the nitrile group and to con-

clude the reaction with alcohol. Thus it was possible to obtain

Card 1/3

SOV/79-29-9-50/76

Synthesis of the Imino Esters and the Esters of &-Cyanovaleric Acid

an almost solid hydrochloric methylimino ester of 6-cyancvaleric acid (I) from adiponitrile, methanol, and HCl in ether-, benzene-, dioxane- or carbon tetrachloride medium in theoretical. yields. (Scheme 1). The molecular compound (II) of the hydrochloride of (I) and of adiponitrile resulted from the same components in a molar ratio of 1:1:0.5. Molecular compounds of this type have hitherto not been described. According to the reaction described in publications (Refs 3.5) i.e. in a molar ratio 1:1:1 and in the passing of hydrogen chloride through the ether or benzene solution of adiponitrile and methanol also the molecular compound (II) was formed besides the hydrochloride of (I). By hydrolyzing the reaction product a mixture of methyl ester of δ-cyanovaleric acid (60% yield) and adiponitrile were formed. However, it was not possible to obtain the methyl ester of δ -cyanovaleric acid (III) from this mixture since it becomes unstable in distillation and regroups into adiponitrile and into the dimethyl esters of adipinic acid; this explains the lack of a description of methyl ester (III) in the patents (Ref 5). By hydrolyzing the hydrochloride of (I) synthesized by the authors, methyl ester (III) could be

Card 2/3

SOV/79-29-9-50/76 Synthesis of the Imino Esters and the Esters of o-Cyanovaleric Acid

obtained which needed no special purification (Table). The ethyl-, n. propyl-, n. butyl, n. amyl-, n. hexyliming esters of δ-cyanovaleric acid were crystallized with difficulty. Therefore they were directly hydrolyzed in the reaction mass under the formation of the corresponding esters of δ-cyanovaleric acid (IV)-(VIII) (Table). There are 1 table and 8 references, 3 of which are Soviet.

SUBMITTED:

July 30, 1958

Card 3/3

KULIKOVA, A.Ye.; ZIL'ERRMAN, Ye.N.; ROGINSKAYA, TS.N.; SMIRMOVA, M.M.

Purification of adiponitrile. Zhur.prikl.khim. 32 no.1:227(NIRA 12:4)

(Adiponitrile)

5(3), 15(9)

SOY/80-32-4-28/47

rath, allo ditabilio kallabe. Dibi dené kabisa dibitera alam

AUTHORS:

Berlin, A.A., Zil'berman, Ye.N., Rybakova, N.A., Sharetskiy, A.M.

and Yancvskiy, D.M.

TITLE:

Investigation of Some Epoxide Stabilizers for Polyvinylchloride (Issledovaniye nekotorykh epoksidnykh stabilizatorov polivinilkhlorida)

PERIODICAL:

Zhurnal prikladnov khimii, 1959, Vol 32, Nr 4, pp 863-868 (USSR)

ABSTRACT:

the of the real drawbacks of ellorine-containing polymers is their how resistance to the effects of heat and light. Variou stabilizers have been proposed for increasing their chermal resistance. The present article furnishes comparative data on the stabilizing effect of some commercial and newly synthesized (by the authors) compounds. The following stabilizers for polyvinylchloride have been synthesized and tested: low-molecular epoxide resins on the base of epichlorohydrin and 2,2-bis(4-cxy-3-methylphenyl)propane, 1,1-bis-(4-oxyphenyl)-cyclohexane, 1,1-bis-(4-oxy-3methylphenyl)-cyclohexane and 2,2-bis-(4-oxy-3-nitrophenyl)propane, cis-9,10-epoxybutyl stearate, epoxidized castor oil and

Card 1/2

sperm oil. It has been shown that these compounds, with exception

SOV/80-32-4-28/47

मेर हो प्रियं । येज्ये जी प्रियं जिल्हा प्रियं है, विशेष्य संक्ष्ये की कि क्षिप्रकार कि अधिक हो का विश्व कि

Investigation of Some Epoxide Stabilizers for Polyvinylchloride

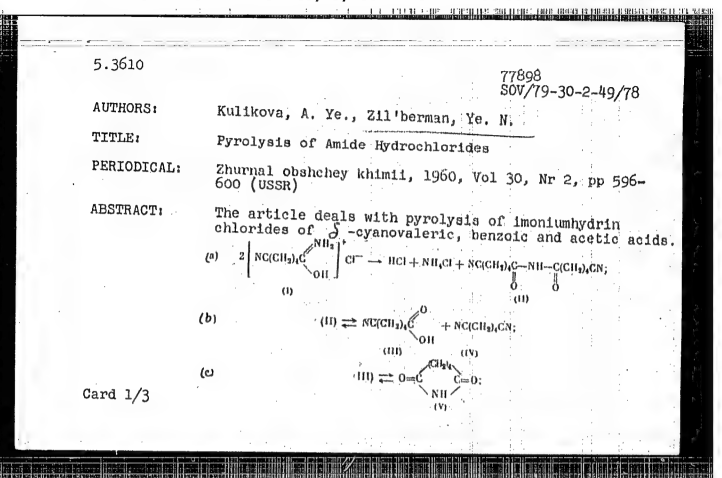
of 2,2-bis-(4-oxy-3-nitrophenyl)-propane are effective thermostabilizers for polyvinylchloride, which improve also physico-mechanical properties of the masticated rubber. The application of mixtures of low-mclecular epoxide resins or epoxidized triglycerides with lead silicate makes it possible to attain a greater thermal resistance of polyvinylchloride and a better quality of the masticated rubber, than the separate application of those stabilizers.

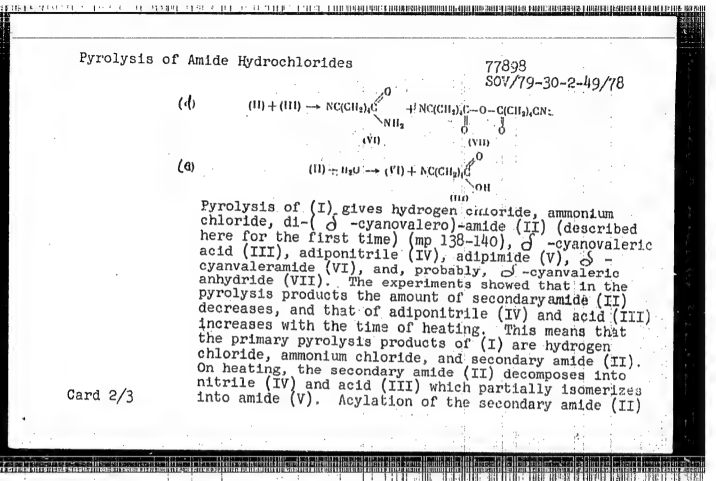
There are 2 graphs, 2 tables and 10 references, 1 of which is Soviet, 5 English, 2 American and 2 German.

SUBMITTED: Set

September 20, 1957

Card 2/2





Pyrolysis of Amide Hydrochlorides

77898 507/79-30-2-49/78

with o -cyanovaleric acid (III) results in an amide (VI) and an anhydride (VII). Pyrolysis of imoniumhydrin chlorides of benzoic and acetic acids was used to confirm the proposed reaction course. The experiments showed that the reactions follow the above rules, but form acid chlorides instead of anhydrides, which can be explained by the hydrogen chloride reaction with either anhydride or secondary amide. It should be noted that thermal decomposition of amide hydrochlorides gives the same products as the pyrolysis of free amides, but the latter occurs at much higher temperatures. There are 2 tables; and 12 references, 2 German, 3 Soviet, 1 Dutch, 4 U.S., 2 U.K. The U.S. and U.K. references are: C. D. Hurd, M. F. Dull, J. Am. Chem. Soc., 54, 2532 (1932); D. Davidson, H. Skovronek, J. Am. Chem. Soc., 80, 376 (1958); D. P. N. Satschell, Chem. and Ind., 1442, (1958); A. W. Ralston, H. J. Harwood, W. O. Pool, J. Am. Chem. Soc., 59, 986 (1937); D. Davidson, M. Karten, J. Am. Chem. Soc., 78, 1066 (1956). February 13, 1959

SUBMITTED:

Card 3/3

CIA-RDP86-00513R002065120017-7 "APPROVED FOR RELEASE: 09/19/2001

78302 5.3610 SOV/79-30-3-56/69

Svetozarskiy, S. V., Razuvayeva, G. A., Zil'berman, AUTHORS:

Ye. N.

Synthesis of 2-Substituted of 4-Pentamethylene-5,6-TITLE:

-Tetramethylene-2,3,4,5-Tetrahydropyrimidines

Zhurnal obshchey khimii, 1960, Vol 30, Nr 3, pp PERIODICAL:

1020-1023 (USSR)

ABSTRACT:

It was shown that the previously obtained (by the authors (ZhOKh, 26, 601, 1956)) 2,4-dipentamethylene-5,6-tetramethylene-2,3,4,5-tetrahydropyrimidine (Ia) can also be prepared by condensation of 2-(1-aminocyclohexyl)-cyclohexanone (II) with ammonia.

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Synthesis of 2-Substituted of 4-Pentamethylene-5,6-Tetramethylene-2,3,4,5-Tetrahydropyrimidines

The following new compounds were obtained by condensation of II and ammonia with different carbonyl compounds. 2,2-Dimethyl-4-pentamethylene-5,6-tetramethylene-2,3,4,5-tetrahydropyrimidine (Ib) was obtained by condensation of II, ammonia, and acctone

(91%), d_h²⁰ 1.002, n_D²⁰ 1.5128; 2-methyl-2-ethyl-4-pentamethylene-5,6-tetramethylene-2,3,4,5-tetrahydro-pyrimidine (Ic) (90.5%), by condensation of 11 am monia,

Card 2/3

 Synthesis of 2-Substituted of 4-Pentamethylene-5,6-Tetramethylene-2,3,4,5-Tetrahydropyrimidines

78302 80V/79-39-3-56/69

and methyl ethyl ketone, d_4^{20} 1.001, n_D^{20} 1.5138; 2-methyl-4-pentamethylene-5,6,-tetramethylene-2,3,4,5-tetrahydropyrimidine (Id) (91%), by condensation of

II, ammonia, and acetaldehyde, d_{\parallel}^{20} 1.02, n_{D}^{20} 1.5202. It was shown that (1) ammonia readily adds to cyclohexanone to form a stable (at low temperatures) compound with a 1:1 molar ratio; (2) this compound is not an intermediate product in the preparation of II and Ia from cyclohexanone and ammonia. There are 7 references, 1 U.K., 4 German, 2 Soviet. The U.K. reference is: R. B. Bradbury, N. C. Hancox, H. H. Hatt, J. Chem. Soc., 1947.

SUBMITTED:

January 22, 1959

Card 3/3

FREYDLIN, L.Kh.; SLADKOVA, T.A.; KUDRYAVTSEV, G.I.; SHEYN, T.I.; AIL'BERMAN, Ye.N.; FEDOROVA, R.G.

Catalytic hydrogenation of aromatic mitriles and the properties of polyamides obtained from p-(9.9° diaminodiethylbenzene). Izv. AN SSSR. Otd.khim.nauk no.9:1715-1715 S '61. (MIRA 14:9)

1. Institut organicheskoy khimii im. N.D.Zelinskogo AN SSSR i Vsesoyuznyy nauchno-issledovatel'skiy institut iskusstvennogo volokna. (Nitriles) (Hydrogenation) (Polyamides)

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ACCESSION NR: AP4040541

8/0064/64/000/006/0408/0416

AUTHOR: Zil!berman, Ye. N.

TITIE: Production of monomers having a high degree of purity for polyemide resins

SOURCE: Khimicheskaya promy*shlennost', no. 6, 1964, 408-416

TOPIC TAGS: literature survey, polyamide resin, polyamide resin monomer, synthesis, caprolactam, hexamethylenediamine sdipate, production, purity, side reactions, Beckmann rearrangement, purification

ABSTRACT: In this survey the domestic and foreign literature on the impurities in monomers for polyamide resins, and the factors causing these impurities, is reviewed. The survey was directed to the synthesis of the two monomers a -caprolactam (made primarily by forming the oxime of cyclohexanone and subsequent Beckmann rearrangement) and hexamethylenediamine adipate (especially the oxidation of cyclohexano or cyclohexanol to edipic acid, preparation of adiponitrile and hydrogenation to hexamethylenediamine, and subsequent reaction with adipic acid). Three groups of side reactions normally accompanying the basic reactions were studied—those resulting from deviations from established technological conditions, or from

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ACCESSION NR: AP4040541

inadequacies in apparatus design for the processes, or those depending on the presence of impurities in the initial products. The necessary measures for suppressing side reactions and removing impurities from these monomers were sought. It was concluded that all the factors affecting the quality of caprolactam and hexamethylendiamine adipate monomers are not known. To obtain high quality monomers it is necessary to suppress side reactions in all (not just the final) stages of production and to separate impurities from the initial and intermediate products in the synthesis of the monomer. Orig. art. has: 26 equations and 9 formulas.

ASSOCIATION: None

SUBMITTED: 00

ENCL: 00

SUB CODE: OC

NO REF SOV: 038

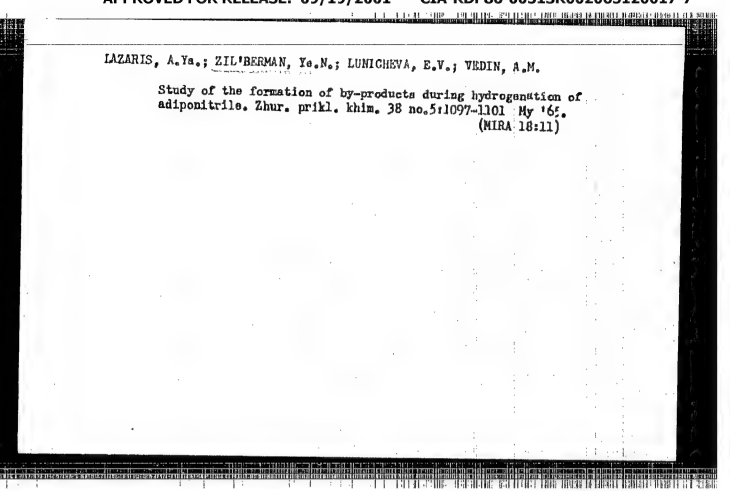
OTHER: 026

Card 2/2

GANINA, V.I.; IVCHER, T.S.; FOMERANTSEVA, E.G.; PEREPLETCHIKOVA, Ye.M.;

ZIL'BERMAN, Ye.N.

Polarographic and spectrophotometric determination of A.—ungaturated ketones in cyclohexanone. Zav. lab. 30
no.5:541-542 '64. (MIRA 17:5)



L 44348-66 EWT(m)/EWP(j)/T IJP(c) WW/RM SOURCE CODE: UR/0191/66/000/004/0003/0004 ACC NR: AP6023056 (A)AUTHOR: Zegel'man, V. I.; Zil'berman, Ye. N.; Kotlyar, I. B.; Svetozarskiy, S. V. omare design a facility of the contraction of the c 29 ORG: none B: TITLE: Low temperature/emulsion polymerization of vinyl chloride SOURCE: Plasticheskiye massy, no. 4, 1966, 3-4 TOPIC TAGS: emulsion polymerization, vinyl chloride, polyvinyl chloride, vinyl plastic ABSTRACT: Kinetics of vinyl chloride polymerization was studied at -20°C, pH=3-13, duration 0-4 hours, using a mixture of sodium alkylsulfonates with 14-18 carbon atoms as emulsifier and ammonium persulfate- ferrous sulfate (0-2 g/l (NH4)2S2O8) redox system as initiator. A maximum of 80-85% yields of polyvinyl chloride were obtained with an equimolar ratio of the components of the redox system at pH=3, polymerization duration equal to 2-4 hours, and 1-2% emulsifier. The polymer molecular weight was found to increase with increasing amount of emulsifier used. The low temperature polymerization used in this work gave PVC with 95-100°C glass point. Orig. art. has: 4 13 figures. OTH REF: 001 ORIG REF: 007/ SUBH DATE: none/ SUB CODE: 07/ UDC: 678.743.22 : 66.095.262.3 Card 1/1 blg

ZIL'BERMAN, Ye.N.; STRIZHAKOV, O.D.; PEREPLETCHIKOVA, Ye.M.

Studying the thermal decomposition of the ester plasticizers of polyvinyl charide. Plast. massy no. 12:29-32 *65 (MIRA 19:1))

<u> na natura de la compaño de la seguida de la compaña de l</u>

PYRYALOVA, P.S.; ZIL'EERMAN, Ye.N.

Reduction of eromatic and branched aliphatic mitriles with stannous chloride. Izv. vys. ucheb. zav.; khim. 1 khim. tekh.
8 no.1;82-8? '65. (MIRA 18:6)

1. Gor'kovskiy politekhnicheskiy institut imeni Zhdanova, kafedra tekhnologii organicheskogo sintesa.

